

BABYNET Covered Diagnoses

Children with documentation that any condition on this list has been professionally diagnosed are eligible for BabyNet services based on “established risk”.

10p13 Deletion	Cleft hands bilertal
11q Deletion	Coffin Lowry syndrome
13q Syndrome	Cornelia de Lange
18q Deletion Syndrome	Cortical blindness
49xxxxxy syndrome	Cri du chat
4p Minus Syndrome	Cystinosis
6q Minus Syndrome	Dandy walker malformation
7q Minus Syndrome	Down syndrome (trisomy 21)
Agenesis of the corpus callosum	Duplication short arm chromosome #20
Albinism	Encephalocele
Amniotic band syndrome	Fazio-Londe disease
Amyoplasia congenita disrutice sequence	Fetal alcohol syndrome
Anencephaly	Fragile X
Angelman syndrome	Glaucoma w/visual impairment
Anophthalmia	Grade IV intraventricular hemorrhage (IVH)
Arginosuccinic aciduria	Hearing loss ≥ 20 db
Arthrogryposis	Hemiparesis
Asphyxia	Herpes encephalitis
Athetoid CP	Holoprosencephaly
Auditory neuropathy	Hydranencephaly
Autism spectrum disorders (ASD)	Hydrocephaly
Automatic eligibility nos	Incontinentia pigmenti syndrome
Bilateral micromelia	Infantile spasms
Bilateral optic nerve coloboma	Isochrome 18 p syndrome
Bilateral retinal detachment w/blindness	Kabuki syndrome
Bilateral visual acuity $\leq 20/70$ corrected vision best eye	Karsch-neugebauer syndrome
Birthweight ≤ 1200 grams	Klinefelter syndrome
Carpenter's syndrome	Krabbe's disease
Cataracts w/ visual impairment	Larsen syndrome
Caudal regression syndrome	Lebers amaurosis
Cerebral palsy (CP)/static encephalopathy	Lennox-gastaut syndrome
Charge association	Lissencephaly syndrome
Citrulinemia	Lowe syndrome (oculo-cerebro-renal)

Marshal Smith Syndrome
Melnick-Frazier
Microdactyly
Midas syndrome
Miller-dieker syndrome
Mobius sequence
Mosaic trisomy 8
MPS (mucopolysaccharidosis)
MSUD (maple syrup urine disease)
Myelodysplasia
Myotonic dystrophy
Myotubular myopathy
Neural tube defects
Opitz syndrome
Optic nerve atrophy
Ornithine-carbamyl-transferase deficiency
Osteogenesis imperfecta
Pachygyria
Pallister-killian syndrome
Pathologic head growth
Perinatal asphyxia, severe
Pervasive developmental disorder (ASD)
Phocomelia
PKU
Prader-willi syndrome
Prematurity (gestational age ≤ 28 wks)
Propionic acidemia
R.O.P. stage 5 & 6 retrolental fibroplasia
Retinitis pigmentosa
Retinoblastoma
Rhizomelic chondrodysplasia punctata
Ring chromosome 13
Schizencephaly
Seckel syndrome
Seizures w/congenital brain malformation
Septo-optic dysplasia
Severe attachment disorder (ASD)
Spastic diplegia

Spastic hemiplegia
Spastic quadriplegia
Spina bifida
Spinal cord injury
Spinal muscular atrophy
Stickler syndrome
Syringohydromyelia
Tar syndrome
Tay-sachs disease
Tetrasomy 12p
Trisomy 1
Trisomy 10
Trisomy 13
Trisomy 18
Trisomy 4
Trisomy 8
Trisomy 9
Tuberous sclerosis
Turner's syndrome
Vater syndrome, with limb anomalies
Velo-cardio-facial syndrome
Waardenberg syndrome
Werdnig-hoffman
William's syndrome
Wolfe-Hershorn syndrome